

Spectroscopic Investigation of Atmospheric Pressure Counterflow Diffusion Flames Inhibited by Halons

Kevin L. McNesby Robert G. Daniel Jeffrey M. Widder Andrzej W. Miziolek

ARL-TR-857

September 1995



APPROVED FOR PUBLIC RELEASE; DISTRIBUTION IS UNLIMITED.

19951012 001

DTIC QUALITY INSPECTED 5

NOTICES

Destroy this report when it is no longer needed. DO NOT return it to the originator.

Additional copies of this report may be obtained from the National Technical Information Service, U.S. Department of Commerce, 5285 Port Royal Road, Springfield, VA 22161.

The findings of this report are not to be construed as an official Department of the Army position, unless so designated by other authorized documents.

The use of trade names or manufacturers' names in this report does not constitute indorsement of any commercial product.

REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Hinghaya, Suite 1204, Artington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0188). Washington, DC 20503.

Davis Highway, Suite 1204, Arlington, VA 22202	-4302, and to the Office of Management and	Budget, Paperwork Reduction Proje	ect (0704-0188), Washington, DC 20503.		
1. AGENCY USE ONLY (Leave blan		3. REPORT TYPE AND			
	September 1995	Final, May 94–A			
4. TITLE AND SUBTITLE			5. FUNDING NUMBERS		
Spectroscopic Investigation of A	Atmospheric Pressure Counterf	low Diffusion Flames	PR: 1L161102AH43		
Inhibited by Halons			PR: ILIOI102AH43		
6. AUTHOR(S)					
Kevin L. McNesby, Robert G.	Daniel, Jeffrey M. Widder, ar	nd			
Andrzej W. Miziolek					
7. PERFORMING ORGANIZATION NA	ME(S) AND ADDRESS(ES)		8. PERFORMING ORGANIZATION REPORT NUMBER		
U.S. Army Research Laborator ATTN: AMSRL-WT-PC	У		ARL-TR-857		
Aberdeen Proving Ground, MI	21005-5066		7HCE-1K-057		
Abeldeen Hoving Glound, Mi	7 21003-3000				
9. SPONSORING/MONITORING AGE	NCY NAME(S) AND ADDRESS(ES	5)	10. SPONSORING / MONITORING		
			AGENCY REPORT NUMBER		
11. SUPPLEMENTARY NOTES					
11. 3011 ZEMZM: ANT NO 123					
12a. DISTRIBUTION / AVAILABILITY S	TATEMENT		12b. DISTRIBUTION CODE		
Approved for public release; distribution is unlimited.					
r,					
13. ABSTRACT (Maximum 200 words)	······································			
Infrared spectra of atmosphe	ric pressure counterflow diffu	sion flames inhibited by	Halons and a few of their potenti	al	
			to spectra of similar flame system		
			ion methane/air flames inhibited b		
CF ₃ Br, CF ₂ H ₂ , and CF ₄ , the two	major fluorine-containing cor	nbustion products are HF	and CF ₂ O. A correlation is show	vn	
			interflow diffusion flames inhibite		
			to be the only fluorine-containing		
			extinguishing atmospheric pressu		
			flame inhibitant testing over a wic		
			pressure inhibition of real fires b		
	Halons, CF ₂ O may be a good indicator of inhibitor efficiency when that inhibition is at least partly accomplished by chemical scavenging of reactive combustion intermediates.				
chemical scavenging of feactive combustion intermediates.					
14. SUBJECT TERMS			15. NUMBER OF PAGES		
flame inhibition, halon replacement, Fourier transform infrared (FT-IR) spectroscopy					
diffusion flames			16. PRICE CODE		
	8. SECURITY CLASSIFICATION	19. SECURITY CLASSIFIC	ATION 20. LIMITATION OF ABSTR	ACT	
OF REPORT UNCLASSIFIED	OF THIS PAGE UNCLASSIFIED	OF ABSTRACT UNCLASSIFIED	SAR		
OLICEMOSITIED	OT 1CEU00ILIED	OTTOPURATION.	אטעי		

ACKNOWLEDGMENT

The authors wish to acknowledge the support of the Strategic Environmental Research and Development Program (SERDP) of the U.S. Department of Defense (DOD). We would also like to thank Dr. Anthony Hamins of NIST for the loan of the counterflow diffusion burner.

Accesi	ọn For		
DTIC	ounced	X	
By Distribution /			
Availability Codes			
Dist	Avail a Spec		
A-1			

TABLE OF CONTENTS

		Page
	ACKNOWLEDGMENT	iii
	LIST OF FIGURES	vii
1.	INTRODUCTION	1
2.	EXPERIMENTAL	2
2.1 2.2 2.3	Atmospheric Pressure Counterflow Diffusion Burner	2 3 3
3.	RESULTS AND DISCUSSION	5
4.	CONCLUSION	10
5.	REFERENCES	13
	DISTRIBUTION LIST	15

LIST OF FIGURES

<u>Figure</u>		Page
1.	A schematic of the atmospheric pressure counterflow diffusion burner used in these experiments	3
2.	A schematic of the low-pressure burner apparatus used in these experiments, showing the burner inside the evacuable chamber	4
3.	The FT-IR absorbance spectrum measured through an atmospheric pressure counterflow diffusion methane/air flame	6
4.	The FT-IR absorbance spectrum measured through an atmospheric pressure counterflow diffusion methane/air flame doped with 1.0% CF ₃ Br	6
5.	The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with successively increasing amounts of CF ₃ Br. Note the increase in the CF ₂ O feature near 1,950 cm ⁻¹	7
6.	The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with CF ₃ Br, CF ₂ H ₂ , and CF ₄	7
7.	The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with 1.0% CF ₄ and an atmospheric pressure counterflow diffusion methane/oxygen flame doped with 1.0% CF ₄	8
8.	The FT-IR absorbance spectra measured through a low-pressure methane/oxygen flame with and without 2.6% CF ₃ Br added to the premixture	9
9.	The FT-IR absorbance spectrum measured through a low-pressure methane/oxygen flame with 15% CF ₃ Br added to the premixture	9

1. INTRODUCTION

Because of concern over the depletion of stratospheric ozone, production and sale of the widely used Halon 1301 (CF₃Br) and Halon 1211 (CF₂C1Br) have been banned (Copenhagen 1987) as of 1 January 1994. A search is presently underway for suitable replacements. To be an acceptable replacement, the new inhibitant must have high fire suppression efficiency, low toxicity, zero residue, compact storage capability, rapid dispersion upon release, and high materials and systems compatibility (Philipczak 1993). The need for environmentally friendly alternatives equal to, or surpassing, the flame inhibition efficiency of Halons 1301 and 1211 is especially important for critical applications encountered in the military. Scenarios range from extinguishment of electrical fires in computer facilities to suppression of a mist-fireball explosion in an armored vehicle following penetration of the fuel cell by a projectile (Finnerty and Polyanski 1993).

The overall goal of the Halon (a contraction of <u>hal</u>ogenated hydrocarbon) alternatives work being conducted at the U.S. Army Research Laboratory (ARL) is the experimental validation of halogen flame mechanisms developed at the National Institute of Standards and Technology (NIST) (Burgess et al. 1994). Once validated, these models will be used to predict inhibitor efficiency and toxic gas by-product formation. The experimental approach employed at ARL uses infrared tunable diode laser (TDL) (Hanson and Falcone 1978) and Fourier transform infrared (FT-IR) spectroscopies to measure in-situ flame temperatures and concentrations of species participating in the combustion occurring in low-pressure premixed and atmospheric pressure counterflow diffusion flames doped with small amounts of Halons and candidate Halon alternatives. Vibrational spectroscopy was chosen as the diagnostic technique because the measurement does not perturb the flame, and because nearly all of the combustion participants (with the exception of homonuclear diatomics) may be observed in simple infrared absorption spectra.

The two flame systems were selected for different reasons. Flat, low-pressure laminar flow premixed-gas flames are used because at low pressure, flame regions are expanded. This expansion provides better spatial resolution for probing preheating and combustion flame zones (Biordi, Lazzara, and Papp 1975). Atmospheric pressure counterflow diffusion flames are used because these flames may closely approximate real fire scenarios, where diffusion and nonpremixed combustion are important (Linteris, in press [a]). The experimental protocol involves qualitative flame species measurements using FT-IR spectroscopy, while spectral temperature (Ouyang, Varghese, and Cline 1989) and species concentration flame profiles are determined using tunable diode laser spectroscopy. Results from experiments in our lab using diode laser absorption spectroscopy have been reported elsewhere (Daniel et al. 1994).

A drawback to the use of FT-IR spectroscopy for the investigation of inhibited flames is the limited spatial resolution afforded by the polychromatic probe beam. Although the output beam waist may be apertured to less than 1 mm, this results in significant loss of throughput, which decreases the signal-to-noise ratio in the spectra. This combination makes obtaining spatially resolved information difficult. Still, it should be pointed out that some researchers have had success obtaining spatial resolution through combusting systems using FT-IR spectroscopy (Solomon et al. 1986; McNesby and Fifer 1993). Although no effort is made here to quantify spectra of inhibited flames measured using FT-IR spectroscopy, significant insight into the nature of flame inhibition may be gained from qualitative interpretation of FT-IR spectra. This insight into the nature of the inhibited flame is the subject of this report.

2. EXPERIMENTAL

2.1 Atmospheric Pressure Counterflow Diffusion Burner. The atmospheric pressure counterflow diffusion burner is shown schematically in Figure 1. The burner assembly was fabricated at NIST. A brief explanation of the operation of the apparatus is as follows. Fuel (methane) is flowed at atmospheric pressure into the flame region from below. Oxidizer (oxygen or air) and inhibitant is flowed into the flame region from above. The flame appears as a thin, flat luminous disc (with slight edge curvature pointing up toward the exhaust shroud) located between the fuel and oxidizer ports. Flame position in the volume between fuel and oxidizer ports is determined by gas flow rates and stoichiometry. For neat methane/oxygen flames using equal fuel and oxidizer flow rates, the flame disc is located nearer to the oxidizer port because of the stoichiometry of the methane/oxygen combustion reaction. All gases are exhausted from the flame region through an exhaust port that forms a shroud around the oxidizer port.

For the flames studied using the atmospheric pressure counterflow diffusion burner, typical flow rates were 600 ml/min oxygen and 500 ml/min methane. When air was used as the oxidizer, the air flow rate was 2.2 l/min and the methane flow rate was 1.1 l/min. Inhibitant flow varied up to a maximum of 1.3% of the total flow for each system investigated. These flow parameters were selected because they gave the most stable flame for that particular fuel/oxidizer combination.

Flow was controlled by an MKS Instruments Inc. type 147B gas flow controller. Although the burner exhaust shroud was connected to a high-volume vacuum pump, it was necessary to contain the atmospheric pressure counterflow diffusion burner within a large box equipped with optical ports and a chimney attached to a fume hood. This arrangement was to prevent noxious fumes (HF and CF_2O) from entering the main laboratory.

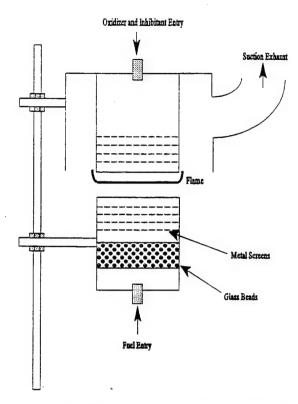


Figure 1. A schematic of the atmospheric pressure counterflow diffusion burner used in these experiments.

- 2.2 Low-Pressure Burner. Most of the low-pressure burner experimental apparatus has been described in a previous publication (Daniel, McNesby, and Miziolek 1993). The low-pressure premixed methane/oxygen flame was supported on a water-cooled, 6-cm-diameter stainless steel fritted, flat flame burner (McKenna Industries). Gases were mixed just prior to entering a final mixing chamber immediately below the fritted burner head. Gas flow was controlled by an MKS type 147B gas flow controller. The low-pressure burner was mounted on a translational stage, which was mounted to a linear motion feedthrough. This low-pressure burner assembly was contained in an evacuable chamber equipped with CaF₂ windows. Pressure was maintained within the chamber by a Heraeus-Leybold Model SV-100 rotary vacuum pump and controlled using an electrically actuated MKS type 253A butterfly valve. Pressure inside the chamber was monitored using MKS type 390 capacitance manometers. Typical flow rates were 200 ml/min oxygen and 100 ml/min methane. Halon flow rates varied up to 15% of the total flow. Typical pressure within the chamber during collection of flame spectra was 20 torr. The experimental apparatus is shown in Figure 2.
- 2.3 <u>FT-IR Spectrometer</u>. The FT-IR spectrometer was manufactured by Mattson Instruments. All counterflow diffusion flame spectra were measured at 4 cm⁻¹ resolution employing triangular apodization

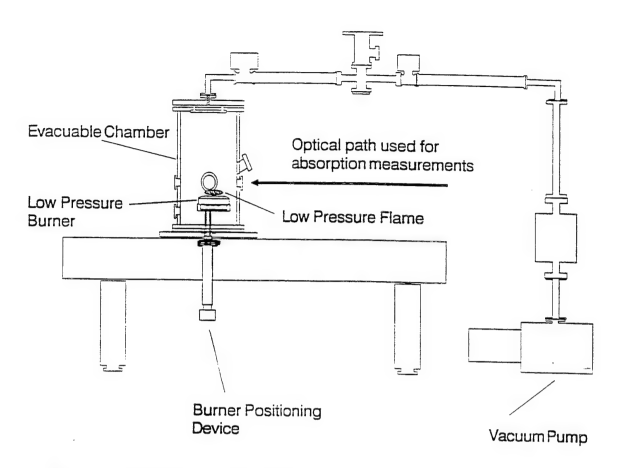


Figure 2. A schematic of the low-pressure burner apparatus used in these experiments, showing the burner inside the evacuable chamber.

using coaddition of 128 scans. Total scan time was approximately 4 min. The infrared beam path was not purged for these experiments. Each single beam flame spectrum was ratioed to a spectrum measured over the same path, but with no flame or fuel/oxidizer/inhibitor mixture present. Since the minimum beam waist diameter of the spectrometer was approximately equal to the distance between fuel and oxidizer ports in the counterflow diffusion burner (see Figure 1), no spatial resolution within the flame was possible, and the resultant spectra show both precursors (fuel and inhibitant) and combustion products.

For the low-pressure experiments, spectra were measured at 1 cm⁻¹ resolution, employing triangular apodization using coaddition of 500 scans. Total scan time was approximately 20 min. The infrared probe beam path was purged with dry nitrogen, and each single beam flame spectrum ratioed to a spectrum through the purged beam path with no flame present and the low-pressure chamber evacuated. The probe beam aperture was set to a diameter of 4 mm when passing through the low-pressure chamber. Since the beam position of the infrared probe was fixed, spectra of different lines-of-sight through the low-pressure flame were measured by moving the burner assembly within the evacuable chamber.

For each set of experiments, reflection of infrared emission through the spectrometer and back onto the detector was checked by collecting spectra with the tungsten carbide filament turned off. In all cases, no interferometer-modulated emission was observed. All gases used in these experiments were of spectral purity and were obtained from Matheson, Inc., except for the CF₃Br, which was obtained from PCR Inc. All gases were used as supplied without further purification.

3. RESULTS AND DISCUSSION

Figure 3 shows the absorbance spectrum through an atmospheric pressure counterflow diffusion methane/air flame with no agent added. The methane fuel is the most prominent feature in the spectrum. Figure 4 shows the same flame doped through the oxidizer port side with 1.0% CF₃Br. Addition of CF₃Br beyond 1% of the total flow causes flame flicker and eventual extinguishment, in agreement with previous studies of diffusion flame extinguishment by CF₃Br (Bajpai 1974). Interesting features of this spectrum are the presence of carbonyl difluoride (CF₂O), near 1,950 cm⁻¹, hydrogen fluoride (HF), near 3,700 cm⁻¹, and Halon 1301 (CF₃Br), near 1,200 cm⁻¹. Both HF and CF₂O are highly toxic and corrosive gases (Sheinson, Musick, and Carhart 1981); therefore, care must be exercised when doping any flame with Halon 1301 when extinguishment is either not a goal or does not occur immediately. Figure 5 shows spectra of the atmospheric pressure methane/air counterflow diffusion flame with successive additions of Halon 1301. It is evident from these spectra that CF₂O formation in these flames is dependent upon the Halon 1301 dopant level.

Figure 6 shows spectra of atmospheric pressure counterflow diffusion methane/air flames doped at the 1% level with Halon 1301 (CF_3Br), 1.3% CF_4 , and 1.3% CF_2H_2 . For the CF_2H_2 doped flame, there is a considerable amount of HF formation near 3,700 cm⁻¹. For the CF_4 doped flame, there is no indication of HF or CF_2O formation, presumably because the CF bond in CF_4 is not broken in the flame (Linteris, in press [b]). The degree to which CF_2O is formed in the atmospheric pressure methane/air counterflow diffusion flames doped with CF_3Br , CF_2H_2 , and CF_4 parallels their efficiency as flame inhibitors determined using cup burner experiments ($CF_3Br > CF_3H > CF_2H_2 > CF_4$) (Linteris, in press [b]).

Figure 7 shows spectra of atmospheric pressure methane/air and methane/oxygen counterflow diffusion flames doped with CF₄. The hotter methane/oxygen flame shows significant HF formation, indicating cleavage of the C-F bond in CF₄. It should be noted that there is no spectral evidence of CF₂O formation in this flame, even though HF is formed. For this reason, we believe that it is questionable to correlate HF formation with inhibitor efficiency.

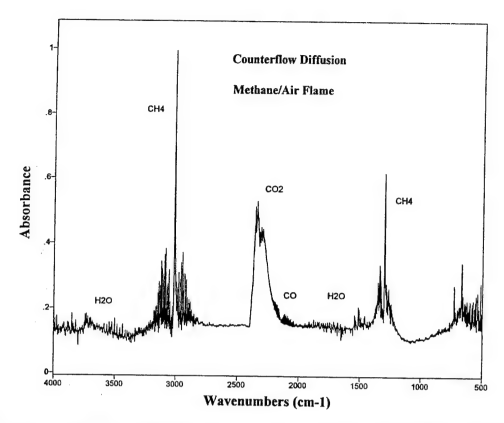


Figure 3. The FT-IR absorbance spectrum measured through an atmospheric pressure counterflow diffusion methane/air flame.

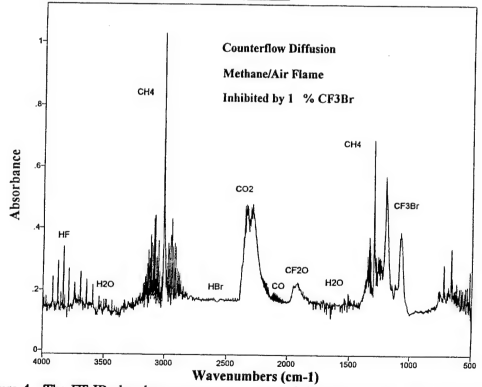


Figure 4. The FT-IR absorbance spectrum measured through an atmospheric pressure counterflow diffusion methane/air flame doped with 1.0% CF₃Br.

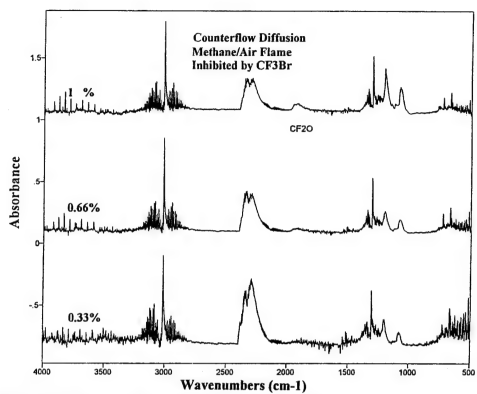


Figure 5. The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with successively increasing amounts of CF₃Br. Note the increase in the CF₂O feature near 1950 cm⁻¹.

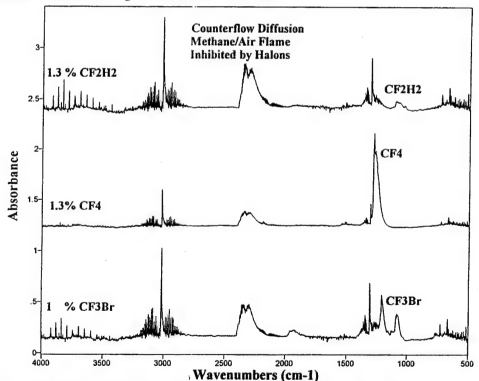


Figure 6. The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with CF₂Br, CF₂H₂, and CF₄.

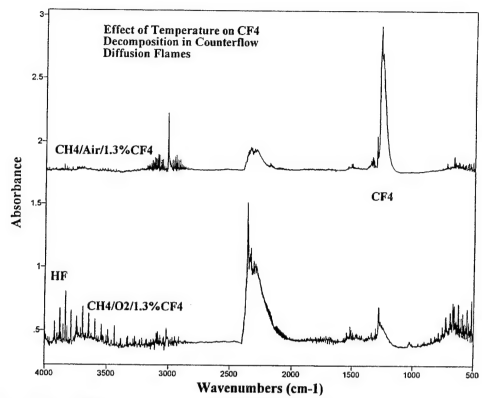


Figure 7. The FT-IR absorbance spectra measured through an atmospheric pressure counterflow diffusion methane/air flame doped with 1.0% CF₄ and an atmospheric pressure counterflow diffusion methane/oxygen flame doped with 1.0% CF₄.

Figure 8 shows the spectra of a low-pressure premixed-gas stoichiometric methane/oxygen flame with and without addition of 2.6% CF₃Br. In contrast to the counterflow diffusion atmospheric pressure methane/air flame doped with CF₃Br, the low-pressure flame shows no evidence of CF₂O formation near 1,950 cm⁻¹. Figure 9 shows the spectrum of the low-pressure premixed-gas methane/oxygen flame doped with 15% CF₃Br. Evident from this spectrum is copious HF formation near 3,700 cm⁻¹ and significant HBr formation near 2,600 cm⁻¹. However, even at this high CF₃Br dopant level, there is no indication of CF₂Br formation at any probe beam position within the flame.

Tunable diode laser infrared absorption measurements of low-pressure flames inhibited by CF_3Br have shown that CF_2O is formed and then rapidly consumed. Modeling studies of these results indicate that at high temperatures, CF_2O is decomposed before leaving the luminous zone of the flame (Miziolek et al., in press). We believe that with line-of-sight spatial resolution of less than 1 mm and spectral resolution better than 1 cm⁻¹, CF_2O would have been detected in these flames using FT-IR spectroscopy.

It is noteworthy that Halon 1301 was unable to extinguish a low-pressure premixed gas stoichiometric methane/oxygen flame even at percent total flow levels 15 times higher than those required to extinguish

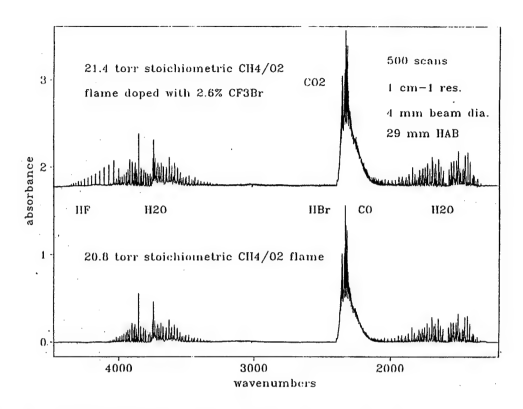


Figure 8. The FT-IR absorbance spectra measured through a low-pressure methane/oxygen flame with and without 2.6% CF₃Br added to the premixture.

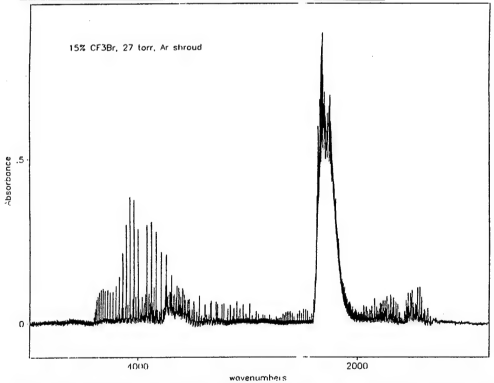


Figure 9. The FT-IR absorbance spectrum measured through a low-pressure methane/oxygen flame with 15% CF₃Br added to the premixture.

atmospheric pressure counterflow diffusion flames. This suggests that the inhibiting effect of Halon 1301 may be most important in flame regions where combustion is nonstoichiometric and diffusion fed. These conditions describe most "real" fires.

Finally, from Figures 4, 8, and 9, it should be noted (assuming similar infrared absorption cross sections for HF and HBr) that there is much less HBr formation than would be expected based on the amounts of HF formation and the molecular formula of Halon 1301 (CF₃Br). This supports calculations that show the rapid removal of HBr from the flame system by reaction (Safieh, Vandooren, and Tiggelin 1982) with H and OH, and the formation of molecular bromine, which is not observable using infrared absorption spectroscopy.

4. CONCLUSION

Although the data are somewhat limited, we believe several conclusions may be reached. Most importantly, we believe the data show that inhibitor effect is highly dependent on the type of combustion system being investigated. Specifically, for atmospheric pressure counterflow diffusion methane/air flames inhibited by CF₃Br, large amounts of CF₂O are observed.

Conversely, for low-pressure premixed-gas stoichiometric laminar methane/oxygen flames inhibited by CF₃Br, no measurable (by FT-IR) CF₂O is observed. This suggests that inhibition of real flames by Halon 1301 may be most important in regions of nonstoichiometric combustion.

Additionally, CF₂O may be a good indicator of the degree of chemical flame inhibition by Halons. Also, it appears that although CF₄ may decompose in hot flames to yield HF, the absence of CF₂O formation indicates that CF₄ does not participate in flame inhibition by the radical scavenging mechanism assumed to be responsible for chemical flame inhibition by Halon 1301. Finally, both low-pressure and atmospheric pressure flames inhibited by Halon 1301 show less HBr formation than would be expected based solely upon the amount of HF observed and the molecular formula of the inhibitant (CF₃Br). This supports the mechanism of rapid HBr removal from the flame via reactions with H and OH, and the conversion of Br atoms to molecular bromine.

We believe that this rather straightforward experimental setup is a valuable tool and may be used to see differences in flame inhibition of real fires by different agents. The technique may be a useful screening test for candidate replacement agents prior to full-scale testing. We are presently engaged in the construction of a new counterflow diffusion burner that will provide larger flame surface area and be capable of low-pressure operation.

5. REFERENCES

- Bajpai, S. N. Journal of Fire and Flammability, vol. 5, p. 255, 1974.
- Biordi, J. C., C. P. Lazzara, and J. F. Papp. "Halogenated Fire Suppressants." ACS Symposium Series 16, American Chemical Society, Washington, DC, 1975.
- Burgess, D., W. Tsang, M. R. Zachariah, and P. R. Westmoreland. "Fluorinated Hydrocarbon Flame Suppression Chemistry." Preprints of papers presented at The American Chemical Society National Meeting (207th), pp. 141–146, San Diego, CA, 13–17 March 1994.
- Copenhagen. Revisions to "The Montreal Protocol on Substances That Deplete the Ozone Layer." 1987.
- Daniel, R. G., K. L. McNesby, and A. W. Miziolek. "A Tunable Diode Laser Study of Premixed Methane/Oxygen Flames Inhibited by Halogenated Compounds." Technical Meeting, Proceedings of the Eastern States Section, The Combustion Institute, pp. 495–498, October 1993.
- Daniel, R. G., K. L. McNesby, A. W. Miziolek, D. R. F. Burgess, Jr., P. R. Westmoreland, W. Tsang, and M. R. Zachariah. "IR Laser Absorption and Modeling Studies of Hydrocarbon Flames Inhibited by Candidate Halon Replacement Compounds." Proceedings of the 1994 Halon Options Technical Working Conference, pp. 229–240, New Mexico Engineering Research Institute, Albuquerque, NM, 1994.
- Finnerty, A. E., and S. Polyanski. "Using Powderpacks for Passive Fire Protection of Military Vehicles." <u>Journal of Fire Science</u>, vol. 2, pp. 242–252, 1993.
- Hanson, R. K., and P. K. Falcone. Applied Optics, vol. 17, pp. 2477-2480, 1978.
- Linteris, G. T. "Acid Gas Production in Inhibited Diffusion Flames." Halon Replacements: Technology and Science, ACS Symposium Series, edited by A. W. Miziolek and W. Tsang, in press (a).
- Linteris, G. T. "The Effect of Inhibitor Concentration on the Inhibition Mechanism of Fluoromethanes in Premixed Methane-Air Flames." ACS Symposium Series, edited by A. W. Miziolek, and W. Tsang, in press (b).
- McNesby, K. L., and R. A. Fifer. "Tomographic Analysis of Line of Sight Infrared Spectra of Low Pressure Flames." ARL-TR-108, U.S. Army Research Laboratory, Aberdeen Proving Ground, MD, April 1993.
- Miziolek, A. W., A. E. Finnerty, R. G. Daniel, K. L. McNesby, W. Tsang, V. I. Babushok, M. R. Zachariach, P. R. Westmoreland, and D. R. F. Burgess, Jr. "Fundamental Studies of Fire Extinguishment for Predicting Halon Alternative Compound Behavior." Proceedings of the 1994 Army Science Conference, in press.
- Ouyang, X., P. L. Varghese, and D. S. Cline. <u>Advances in Laser Science IV</u>. American Institute of Physics, p. 750, edited by J. L. Gole, D. F. Heller, M. Lapp, and W. C. Swalley, New York, 1989.
- Philipczak, R. A. "Relative Extinguishment Effectiveness and Agent Decomposition Products of Halon Alternative Agents." Proceedings of the 1993 Halon Options Technical Working Conference, pp. 149–159, New Mexico Engineering Research Institute, Albuquerque, NM, 1993.

- Safieh, H. Y., J. Vandooren, and P. J. Tiggelin. "Experimental Study of Inhibition Induced by CF₃Br in a CO-H₂-O₂-Ar Flame." Nineteenth Symposium (International) on Combustion, The Combustion Institute, pp. 117–126, 1982.
- Sheinson, R. S., J. K. Musick, and H. W. Carhart. <u>Journal of Fire and Flammability</u>, vol. 12, pp. 229–235, 1981.
- Solomon, P. R., R. M. Carangelo, D. G. Hamblen, and P. E. Best. "Infrared Analysis of Particulates by FTIR Emission/Transmission Spectroscopy." <u>Applied Spectroscopy</u>, vol. 40, pp. 746–759, 1986.

NO. OF COPIES ORGANIZATION

- 2 ADMINISTRATOR
 ATTN DTIC DDA
 DEFENSE TECHNICAL INFO CTR
 CAMERON STATION
 ALEXANDRIA VA 22304-6145
- 1 DIRECTOR
 ATTN AMSRL OP SD TA
 US ARMY RESEARCH LAB
 2800 POWDER MILL RD
 ADELPHI MD 20783-1145
- 3 DIRECTOR
 ATTN AMSRL OP SD TL
 US ARMY RESEARCH LAB
 2800 POWDER MILL RD
 ADELPHI MD 20783-1145
- 1 DIRECTOR
 ATTN AMSRL OP SD TP
 US ARMY RESEARCH LAB
 2800 POWDER MILL RD
 ADELPHI MD 20783-1145

ABERDEEN PROVING GROUND

5 DIR USARL ATTN AMSRL OP AP L (305)

NO. OF NO. OF COPIES ORGANIZATION COPIES ORGANIZATION 1 **HQDA** 2 COMMANDER ATTN SARD TT DR F MILTON US ARMY MISSILE COMMAND PENTAGON ATTN AMSMI RD PR E A R MAYKUT WASHINGTON DC 20310-0103 AMSMI RD PR P R BETTS REDSTONE ARSENAL AL 1 HODA ATTN SARD TT MR J APPEL 1 OFFICE OF NAVAL RESEARCH PENTAGON DEPARTMENT OF THE NAVY **WASHINGTON DC 20310-0103** ATTN R S MILLER CODE 432 800 N OUINCY STREET HODA OASA RDA **ARLINGTON VA 22217** ATTN DR C H CHURCH PENTAGON ROOM 3E486 1 COMMANDER WASHINGTON DC 20310-0103 NAVAL AIR SYSTEMS COMMAND ATTN J RAMNARACE AIR 54111C COMMANDER WASHINGTON DC 20360 US ARMY RESEARCH OFFICE ATTN R GHIRARDELLI 2 **COMMANDER** D MANN NAVAL SURFACE WARFARE CENTER R SINGLETON ATTN R BERNECKER R 13 R SHAW G B WILMOT R 16 P O BOX 12211 SILVER SPRING MD 20903-5000 RSCH TRNGLE PK NC 27709-2211 5 COMMANDER 1 DIRECTOR NAVAL RESEARCH LABORATORY ARMY RESEARCH OFFICE ATTN M C LIN ATTN AMXRO MCS K CLARK J MCDONALD P O BOX 12211 E ORAN RSCH TRNGLE PK NC 27709-2211 J SHNUR R J DOYLE CODE 6110 DIRECTOR 1 WASHINGTON DC 20375 ARMY RESEARCH OFFICE COMMANDER ATTN AMXRO RT IP LIB SERVICES P O BOX 12211 NAVAL WEAPONS CENTER RSCH TRNGLE PK NC 27709-2211 ATTN T BOGGS CODE 388 T PARR CODE 3895 2 **COMMANDER** CHINA LAKE CA 93555-6001 US ARMY ARDEC ATTN SMCAR AEE B D S DOWNS 1 SUPERINTENDENT PCTNY ARSNL NJ 07806-5000 NAVAL POSTGRADUATE SCHOOL **DEPT OF AERONAUTICS** 2 COMMANDER ATTN D W NETZER US ARMY ARDEC **MONTEREY CA 93940** ATTN SMCAR AEE J A LANNON PCTNY ARSNL NJ 07806-5000 3 AL LSCF ATTN R CORLEY **COMMANDER** 1 R GEISLER US ARMY ARDEC J LEVINE ATTN SMCAR AEE BR L HARRIS EDWARDS AFB CA 93523-5000 PCTNY ARSNL NJ 07806-5000

NO. OF		NO. OF	
COPIES	ORGANIZATION	COPIES	ORGANIZATION
1	AFOSR	2	PRINCETON COMBUSTION
	ATTN J M TISHKOFF		RESEARCH LABORATORIES INC
	BOLLING AIR FORCE BASE		ATTN N A MESSINA
	WASHINGTON DC 20332		M SUMMERFIELD
			PRINCETON CORPORATE PLAZA
1	OSD SDIO IST		BLDG IV SUITE 119
	ATIN L CAVENY		11 DEERPARK DRIVE
	PENTAGON		MONMOUTH JUNCTION NJ 08852
	WASHINGTON DC 20301-7100		
		3	DIRECTOR
1	COMMANDANT		SANDIA NATIONAL LABORATORIES
	USAFAS		DIVISION 8354
	ATTN ATSF TSM CN		ATTN S JOHNSTON
	FORT SILL OK 73503-5600		P MATTERN
			D STEPHENSON
1 .	UNIV OF DAYTON RSCH INSTITUTE		LIVERMORE CA 94550
	ATTN D CAMPBELL		PRIORILI CHOLDIO LEUM EDOME
	AL PAP	1	BRIGHAM YOUNG UNIVERSITY
	EDWARDS AFB CA 93523		DEPT OF CHEMICAL ENGINEERING ATTN M W BECKSTEAD
	NAGA		PROVO UT 84058
1	NASA LANGLEY RESEARCH CENTER		PROVO 01 84038
	ATTN G B NORTHAM MS 168	1	CALIFORNIA INSTITUTE OF TECH
	LANGLEY STATION	1	JET PROPULSION LABORATORY
	HAMPTON VA 23365		ATTN L STRAND MS 125 224
	TIANN TON VA 25505		4800 OAK GROVE DRIVE
4	NATIONAL BUREAU OF STANDARDS		PASADENA CA 91109
•	US DEPARTMENT OF COMMERCE		
	ATTN J HASTIE	1	CALIFORNIA INSTITUTE OF TECHNOLOGY
	M JACOX		ATTN F E C CULICK MC 301 46
	T KASHIWAGI		204 KARMAN LAB
	H SEMERJIAN		PASADENA CA 91125
	WASHINGTON DC 20234		
		1	UNIVERSITY OF CALIFORNIA
2	DIRECTOR		LOS ALAMOS SCIENTIFIC LAB
	LAWRENCE LIVERMORE NATIONAL LAB		P O BOX 1663 MAIL STOP B216
	ATTN C WESTBROOK		LOS ALAMOS NM 87545
	W TAO MS L 282		
	P O BOX 808	1	UNIVERSITY OF CALIFORNIA BERKELEY
	LIVERMORE CA 94550		CHEMISTRY DEPARMENT
	DIRECTOR		ATTN C BRADLEY MOORE
1	DIRECTOR		211 LEWIS HALL BERKELEY CA 94720
	LOS ALAMOS NATIONAL LAB ATTN B NICHOLS T7 MS B284		BERNELET CA 94720
	P O BOX 1663	1	UNIVERSITY OF CALIFORNIA SAN DIEGO
	LOS ALAMOS NM 87545		ATTN F A WILLIAMS
	LOS ALANOS INI 07575		AMES B010
			LA JOLLA CA 92093

NO. OF COPIES	ORGANIZATION	NO. OF COPIES	<u>ORGANIZATION</u>
2	UNIV OF CALIFORNIA SANTA BARBARA QUANTUM INSTITUTE ATTN K SCHOFIELD M STEINBERG SANTA BARBARA CA 93106	1	THE JOHNS HOPKINS UNIV CPIA ATTN T W CHRISTIAN 10630 LITTLE PATUXENT PKWY SUITE 202 COLUMBIA MD 21044-3200
	UNIV OF COLORADO AT BOULDER ENGINEERING CENTER ATTN J DAILY CAMPUS BOX 427 BOULDER CO 80309-0427	1	UNIVERSITY OF MICHIGAN GAS DYNAMICS LAB ATTN G M FAETH AEROSPACE ENGINEERING BLDG ANN ARBOR MI 48109-2140
	UNIV OF SOUTHERN CALIFORNIA DEPT OF CHEMISTRY ATTN R BEAUDET S BENSON C WITTIG	1	UNIVERSITY OF MINNESOTA DEPT OF MECHANICAL ENGINEERING ATTN E FLETCHER MINNEAPOLIS MN 55455
1	LOS ANGELES CA 90007 CORNELL UNIVERSITY DEPARTMENT OF CHEMISTRY ATTN T A COOL BAKER LABORATORY ITHACA NY 14853	4	PENNSYLVANIA STATE UNIVERSITY DEPT OF MECHANICAL ENGINEERING ATTN K KUO M MICCI S THYNELL V YANG UNIVERSITY PARK PA 16802
	UNIVERSITY OF DELAWARE CHEMISTRY DEPARTMENT ATTN T BRILL NEWARK DE 19711	2	PRINCETON UNIVERSITY FORRESTAL CAMPUS LIBRARY ATTN K BREZINSKY I GLASSMAN P O BOX 710
	UNIVERSITY OF FLORIDA DEPT OF CHEMISTRY ATTN J WINEFORDNER GAINESVILLE FL 32611		PRINCETON NJ 08540 PURDUE UNIVERSITY SCHL OF AERONAUTICS & ASTRONAUTICS ATTN J R OSBORN
;	GEORGIA INSTITUTE OF TECHNOLOGY SCHOOL OF AEROSPACE ENGINEERING ATTN E PRICE W C STRAHLE B T ZINN ATLANTA GA 30332		GRISSOM HALL WEST LAFAYETTE IN 47906 PURDUE UNIVERSITY DEPARTMENT OF CHEMISTRY ATTN E GRANT
]	UNIVERSITY OF ILLINOIS DEPT OF MECH ENG ATTN H KRIER 144MEB 1206 W GREEN ST URBANA IL 61801	2	WEST LAFAYETTE IN 47906 PURDUE UNIVERSITY SCHL OF MECHANICAL ENGNRNG ATTN N M LAURENDEAU S N B MURTHY TSPC CHAFFEE HALL WEST LAFAYETTE IN 47906

NO. OF COPIES	<u>ORGANIZATION</u>	NO. OF COPIES	ORGANIZATION
1	RENSSELAER POLYTECHNIC INST DEPT OF CHEMICAL ENGINEERING ATTN A FONTUN	1	GENERAL APPLIED SCIENCE LABS INC 77 RAYNOR AVENUE RONKONKAMA NY 11779-6649
	TROY NY 12181		
1	STANFORD UNIVERSITY	1	GENERAL ELECTRIC ORDNANCE SYSTEMS ATTN J MANDZY
	DEPT OF MECHANICAL ENGINEERING ATTN R HANSON STANFORD CA 94305		100 PLASTICS AVENUE PITTSFIELD MA 01203
		1	GENERAL MOTORS RSCH LABS
1	UNIVERSITY OF TEXAS DEPT OF CHEMISTRY		PHYSICAL CHEMISTRY DEPARTMENT ATTN T SLOANE
	ATTN W GARDINER AUSTIN TX 78712		WARREN MI 48090-9055
		2	HERCULES INC
1	VA POLYTECH INST AND STATE UNIV ATTN J A SCHETZ		ATTN W B WALKUP E A YOUNT
	BLACKSBURG VA 24061		P O BOX 210
			ROCKET CENTER WV 26726
1	APPLIED COMBUSTION TECHNOLOGY INC		THE OWN FOR THE
	ATTN A M VARNEY P O BOX 607885	1	HERCULES INC ATTN R V CARTWRIGHT
	ORLANDO FL 32860		100 HOWARD BLVD
			KENVIL NJ 07847
2	APPLIED MECHANICS REVIEWS		ATTIAND DECLICACEEN AS INC.
	ASME ATTN R E WHITE & A B WENZEL	1	ALLIANT TECHSYSTEMS INC MARINE SYSTEMS GROUP
	345 E 47TH STREET		ATTN D E BRODEN MS MN50 2000
	NEW YORK NY 10017		600 2ND STREET NE
	THE VIEW ON THE PROPERTY OF CANONICA CO.		HOPKINS MN 55343
1	TEXTRON DEFENSE SYSTEMS ATTN A PATRICK	1	ALLIANT TECHSYSTEMS INC
	2385 REVERE BEACH PARKWAY	1	ATTN R E TOMPKINS
	EVERETT MA 02149-5900		MN 11 2720
	D		600 SECOND ST NORTH
1	BATTELLE TWSTIAC		HOPKINS MN 55343
	505 KING AVENUE	1	IBM CORPORATION
	COLUMBUS OH 43201-2693		RESEARCH DIVISION
			ATTN A C TAM
	COHEN PROFESSIONAL SERVICES ATTN N S COHEN		5600 COTTLE ROAD SAN JOSE CA 95193
	141 CHANNING STREET		SAN JOSE CA 93193
	REDLANDS CA 92373		IIT RESEARCH INSTITUTE
,	EVVON DECEADON & ENGLOS		ATTN R F REMALY
	EXXON RESEARCH & ENG CO ATTN A DEAN		10 WEST 35TH STREET CHICAGO IL 60616
	ROUTE 22E		CINCILO ID WOIV
	ANNANDALE NJ 08801		

NO. OF COPIES	ORGANIZATION	NO. OF COPIES	ORGANIZATION
1	LOCKHEED MISSILES & SPACE CO ATTN GEORGE LO 3251 HANOVER STREET DEPT 52 35 B204 2	1	SVERDRUP TECHNOLOGY INC ATTN J DEUR 2001 AEROSPACE PARKWAY BROOK PARK OH 44142
	PALO ALTO CA 94304	3	THIOKOL CORPORATION
1	OLIN ORDNANCE ATTN V MCDONALD LIBRARY P O BOX 222 ST MARKS FL 32355-0222		ELKTON DIVISION ATTN R BIDDLE R WILLER TECH LIB
1	PAUL GOUGH ASSOCIATES INC ATTN P S GOUGH		P O BOX 241 ELKTON MD 21921
,	1048 SOUTH STREET PORTSMOUTH NH 03801-5423	3	THIOKOL CORPORATION WASATCH DIVISION ATTN S J BENNETT
1	HUGHES AIRCRAFT COMPANY ATTN T E WARD 8433 FALLBROOK AVENUE		P O BOX 524 BRIGHAM CITY UT 84302
1	CANOGA PARK CA 91303 ROCKWELL INTERNATIONAL CORP	1	UNITED TECHNOLOGIES RSCH CENTER ATTN A C ECKBRETH
1	ROCKETDYNE DIVISION ATTN J E FLANAGAN HB02 6633 CANOGA AVENUE CANOGA PARK CA 91304	1	EAST HARTFORD CT 06108 UNITED TECHNOLOGIES CORP CHEMICAL SYSTEMS DIVISION ATTN R R MILLER
1	SCIENCE APPLICATIONS INC ATTN R B EDELMAN		P O BOX 49028 SAN JOSE CA 95161-9028
	23146 CUMORAH CREST WOODLAND HILLS CA 91364	1	UNIVERSAL PROPULSION COMPANY ATTN H J MCSPADDEN 25401 NORTH CENTRAL AVENUE
3	SRI INTERNATIONAL ATTN G SMITH		PHOENIX AZ 85027-7837
	D CROSLEY D GOLDEN 333 RAVENSWOOD AVENUE MENLO PARK CA 94025	1	VERITAY TECHNOLOGY INC ATTN E B FISHER 4845 MILLERSPORT HIGHWAY EAST AMHERST NY 14051-0305
	STEVENS INSTITUTE OF TECH DAVIDSON LABORATORY ATTN R MCALEVY III HOBOKEN NJ 07030	1	FREEDMAN ASSOCIATES ATTN E FREEDMAN 2411 DIANA ROAD BALTIMORE MD 21209-1525
	SVERDRUP TECHNOLOGY INC LERC GROUP ATTN R J LOCKE MS SVR 2 2001 AEROSPACE PARKWAY BROOK PARK OH 44142		ALLIANT TECHSYSTEMS ATTN C CANDLAND L OSGOOD R BECKER M SWENSON 600 SECOND ST NE HOPKINS MN 55343

NO. OF

COPIES ORGANIZATION

1 US ARMY BENET LABORATORY
ATTN SAM SOPOK
SMCAR CCB B
WATERVLIET NY 12189

ABERDEEN PROVING GROUND

36 DIR USARL

ATTN: AMSRL-WT-P, A HORST

AMSRL-WT-PC,

R A FIFER

G F ADAMS

W R ANDERSON

R A BEYER

S W BUNTE

C F CHABALOWSKI

K P MCNEILL-BOONSTOPPEL

A COHEN

R CUMPTON

R DANIEL

D DEVYNCK

N F FELL

B E FORCH

J M HEIMERL

A J KOTLAR

M R MANAA

W F MCBRATNEY

K L MCNESBY

S V MEDLIN

M S MILLER

A W MIZIOLEK

S H MODIANO

J B MORRIS

J E NEWBERRY

S A NEWTON

R A PESCE-RODRIGUEZ

B M RICE

R C SAUSA

M A SCHROEDER

J A VANDERHOFF

M WENSING

A WHREN

J M WIDDER

C WILLIAMSON

AMSRL-CI-CA, R PATEL

USER EVALUATION SHEET/CHANGE OF ADDRESS

This Laboratory undertakes a continuing effort to improve the quality of the reports it publishes. Your comments/answers to the items/questions below will aid us in our efforts. 1. ARL Report Number <u>ARL-TR-857</u> Date of Report <u>September 1995</u> 2. Date Report Received _____ 3. Does this report satisfy a need? (Comment on purpose, related project, or other area of interest for which the report will be used.) 4. Specifically, how is the report being used? (Information source, design data, procedure, source of ideas, etc.) 5. Has the information in this report led to any quantitative savings as far as man-hours or dollars saved, operating costs avoided, or efficiencies achieved, etc? If so, please elaborate. 6. General Comments. What do you think should be changed to improve future reports? (Indicate changes to organization, technical content, format, etc.) Organization Name **CURRENT** ADDRESS Street or P.O. Box No. City, State, Zip Code 7. If indicating a Change of Address or Address Correction, please provide the Current or Correct address above and the Old or Incorrect address below. Organization OLD Name **ADDRESS** Street or P.O. Box No. City, State, Zip Code (Remove this sheet, fold as indicated, tape closed, and mail.)

(DO NOT STAPLE)